STIC Biotechnology Systems Branch

## RAW SEQUENCE LISTING % ERROR REPORT

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Application Serial Number: /0/573,/30A

Source: /F/0

Date Processed by STIC: 8/29/06

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FOR CRF SUBMISSION AND PATENTIN SOFTWARE QUESTIONS, PLEASE CONTACT MARK SPENCER, TELEPHONE: 571-272-2510; FAX: 571-273-0221

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Revised 01/10/05





**IFWO** 

RAW SEQUENCE LISTING DATE: 08/29/2006
PATENT APPLICATION: US/10/573,130A TIME: 08:46:39

Input Set : A:\Substitute Sequence List-13111-00035-US.txt

Output Set: N:\CRF4\08292006\J573130A.raw

```
3 <110> APPLICANT: Sturmer, Rainer
        Kesseler, Maria
        Hauer, Bernhard
 5
        Friedrich, Thomas
 6
        Breuer, Michael
 9 <120> TITLE OF INVENTION: Methods for the production of
        3-methylamino-1-(thiene-2-yl)-propane-1-ol
10
12 <130> FILE REFERENCE: 13111-00035-US
14 <140> CURRENT APPLICATION NUMBER: US/10/573,130A
                                                               Does Not Comply
15 <141> CURRENT FILING DATE: 2006-03-23
                                                               Corrected Diskette Needed
17 <150> PRIOR APPLICATION NUMBER: PCT/EP2004/010939
18 <151> PRIOR FILING DATE: 2004-09-30
20 <150> PRIOR APPLICATION NUMBER: DE 103 45 772.0
21 <151> PRIOR FILING DATE: 2003-10-01
                                                            ser pp 6,8
23 <160> NUMBER OF SEQ ID NOS: 44
25 <170> SOFTWARE: PatentIn version 3.3
28 <210> SEQ ID NO: 1
30 <211> LENGTH: 47
32 <212> TYPE: PRT
34 <213> ORGANISM: Lactobacillus brevis
37 <400> SEQUENCE: 1
39 Met Ser Asn Arg Leu Asp Gly Lys Val Ala Ile Val Thr Gly Gly Thr
                                                            15
                                       10
40 1
43 Leu Gly Ile Gly Leu Ala Ile Ala Thr Lys Phe Val Glu Glu Gly Ala
                                   25
               20
44
47 Lys Val Met Ile Thr Gly Arg His Ser Asp Val Gly Glu Lys Ala
                               40
48
           35
51 <210> SEQ ID NO: 2
53 <211> LENGTH: 18
55 <212> TYPE: PRT
57 <213> ORGANISM: Candida magnoliae
60 <400> SEQUENCE: 2
62 Ser Asn Ala Leu Val Thr Gly Gly Ser Arg Val Ile Gly Ala Gly Gly
63 1
                   5
66 Phe Ile
70 <210> SEQ ID NO: 3
72 <211> LENGTH: 756
74 <212> TYPE: DNA
76 <213> ORGANISM: Lactobacillus brevis
79 <220> FEATURE:
81 <221> NAME/KEY: CDS
83 <222> LOCATION: (1)..(756)
86 <400> SEQUENCE: 3
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RAW SEQUENCE LISTING DATE: 08/29/2006 PATENT APPLICATION: US/10/573,130A TIME: 08:46:39

Input Set: A:\Substitute Sequence List-13111-00035-US.txt
Output Set: N:\CRF4\08292006\J573130A.raw

87 a	tg t	ct a	ac Asn	cgt Arg	ttg ( Leu <i>i</i>	gat Asp	gga Gly	aaa Lys	gta Val	Ala	atc Ile	gtt Val	aca Thr	Gly	Gly	acg Thr	48
89 1		rat a	at c		5 tta 🤄	act	atc	acc	acq	10 aaq	ttc	qtt	qaa		15 999	gct	96
92 L 93	eu G	ly 1	lle	Gly : 20	Leu	Ala	Ile	Ala	Thr 25	Lys	Phe	Val	Ğlu	Ğlu 30	Gly	Āla	
95 a	ag g	jtc a	atq	att	acc	ggc	cgg	cac	agc	gat	gtt	ggt	gaa	aaa	gca	gct	144
	ys V			Ile	Thr	Gly			Ser	Asp	Val	Gly	Glu 45	Lys	Ala	Ala	
97 99 a	ag a		35 atc	aac	act	cct		40 caq	att	caa	ttt	ttc		cat	gat	tct	192
100	Lys	Ser	Val	Gly	Thr	Pro	Aap	Gln	Ile	Glr	Phe	Phe	Gln	His	Asp	Ser	
101	_	50		_			55					60					
103	tcc	gat	gaa	gac	ggc	tgg	acg	aaa	tta	tto	gat	gca	acg	gaa	aaa	gcc	240
		Asp	Glu	Asp	Gly	Trp	Thr	Lys	Leu	Pne	: Asg 75	) Ala	The	GIU	Lys	Ala 80	
105		aac	cca	att	tet		tta	att	aat	aac		. aaa	ato	aca	att	aac	288
108	Phe	Glv	Pro	Val	Ser	Thr	Leu	Val	Asn	Asn	Ala	Gly	Ile	Ala	Val	Asn	
109		_			85					90					95		
111	aag	agt	gto	gaa	gaa	acc	acg	act	gct	gaa	tgg	gcgt	aaa	cta	tta	gcc	336
112	Lys	Ser	Val			Thr	Thr	Thr			Tr	Arg	Lys			Ala	
113				100					105					110			384
115	gtc	aac	ctt	gat	ggt	gto	Dla:	מלכ	ggt	acc	cga	l tta	ggg	, Tle	Cac	a cgg	204
116	vaı	ASI	115		GIY	vai	Pne	120		1111	. ALC	, ner	125			9	
	ato	aad			ggc	tta	aaa			ato	ato	: aac			tec	atc	432
120	Met	Lvs	Asn	Lvs	Gly	Leu	Gly	Ala	Ser	Ile	: Ile	Asr	Met	Ser	Sei	Ile	
121		130					135					140	)				
123	gaa	ggc	ttt	gtg	ggt	gat	cct	ago	: tta	999	gct	tac	aac	gca	tct	aaa	480
		Gly	Phe	Val	Gly			Ser	Lev	Gly			Asr	ı Ala	. Sez	Lys	
125						150					155		t	+ ~+		160	528
127	999	gcc	gta	cgg	att Tla	Mot	Cor	Twe	Cer	· Als	. gcc	1.01	. gat	CVS	. gcc	cta Leu	320
129	GIY	AIG	val	Arg	165	1400	Jer	Dy 3	, 501	170				. 0,1	175		
	aag	gac	tac	gat		cgg	gta	aac	act	gtt	. cac	: cct	ggc	: tac	ato	aag	576
132	Lys	Āsp	Tyr	Asp	Val	Arg	Val	Asn	Thr	. Val	. His	Pro	Gly	Туг	: Ile	. Lys	
133				180					185					190			
135	aca	cca	ttg	gtt	gat	gac	cta	cca	999	gcc	gaa	gaa	gcg	atg	tca	caa	624
	Thr	Pro			Asp	Asp	Leu			Ala	GIU	1 GIV	205		. sei	Gln	
137	~~~	200	195			a+0		200		aat	· mas	cct			att	gcc	672
140	Ara	Thr	Lvs	Thr	Pro	Met	Glv	His	Ile	Glv	. gat	Pro	Asr	Ast	Ile	Ala	• • •
141	5	210	-,, -				215					220		•			
143	tac	atc	tgt	gtt	tac	ttg	gct	tct	aac	gaa	tct	. aaa	ttt	gca	ace	ggt	720
144	Tyr	Ile	Cys	Val	Tyr	Lev	Ala	Ser	Asn	Glu	Ser	Lys	Phe	: Ala	Thi	Gly	
145						230					235					240	95.5
					gtt												756
148	ser	GTA	rne	val	Val 245	ASP	, сту	GIĀ	TAX	250		GII					
	<21¢	)> <1	EO T	D NO						236	•						
	~~~	1	-× -		-												

RAW SEQUENCE LISTING DATE: 08/29/2006
PATENT APPLICATION: US/10/573,130A TIME: 08:46:39

Input Set: A:\Substitute Sequence List-13111-00035-US.txt
Output Set: N:\CRF4\08292006\J573130A.raw

```
154 <211> LENGTH: 252
156 <212> TYPE: PRT
158 <213> ORGANISM: Lactobacillus brevis
161 <400> SEQUENCE: 4
163 Met Ser Asn Arg Leu Asp Gly Lys Val Ala Ile Val Thr Gly Gly Thr
                                      10
164 1
167 Leu Gly Ile Gly Leu Ala Ile Ala Thr Lys Phe Val Glu Glu Gly Ala
             20
                                                      30
                                  25
171 Lys Val Met Ile Thr Gly Arg His Ser Asp Val Gly Glu Lys Ala Ala
          35
                               40
172
175 Lys Ser Val Gly Thr Pro Asp Gln Ile Gln Phe Phe Gln His Asp Ser
                           55
                                              60
176 50
179 Ser Asp Glu Asp Gly Trp Thr Lys Leu Phe Asp Ala Thr Glu Lys Ala
                                          75
                       70
183 Phe Gly Pro Val Ser Thr Leu Val Asn Asn Ala Gly Ile Ala Val Asn
                                      90
                   85
184
187 Lys Ser Val Glu Glu Thr Thr Thr Ala Glu Trp Arg Lys Leu Leu Ala
                                                     110
                                  105
188
             100
191 Val Asn Leu Asp Gly Val Phe Phe Gly Thr Arg Leu Gly Ile Gln Arg
                                                  125
                              120
195 Met Lys Asn Lys Gly Leu Gly Ala Ser Ile Ile Asn Met Ser Ser Ile
196 130
                           135
                                              140
199 Glu Gly Phe Val Gly Asp Pro Ser Leu Gly Ala Tyr Asn Ala Ser Lys
                                          155
                       150
200 145
203 Gly Ala Val Arg Ile Met Ser Lys Ser Ala Ala Leu Asp Cys Ala Leu
                                      170
                                                          175
                  165
207 Lys Asp Tyr Asp Val Arg Val Asn Thr Val His Pro Gly Tyr Ile Lys
                                                      190
                                   185
              180
208
211 Thr Pro Leu Val Asp Asp Leu Pro Gly Ala Glu Glu Ala Met Ser Gln
212 195
                              200
215 Arg Thr Lys Thr Pro Met Gly His Ile Gly Glu Pro Asn Asp Ile Ala
                                              220
216 210
                           215
219 Tyr Ile Cys Val Tyr Leu Ala Ser Asn Glu Ser Lys Phe Ala Thr Gly
                                                              240
220 225
                       230
                                          235
223 Ser Glu Phe Val Val Asp Gly Gly Tyr Thr Ala Gln
224
                   245
227 <210> SEQ ID NO: 5
229 <211> LENGTH: 472
231 <212> TYPE: DNA
233 <213> ORGANISM: Candida magnoliae
236 <220> FEATURE:
238 <221> NAME/KEY: CDS
240 <222> LOCATION: (1)..(471)
243 <400> SEQUENCE: 5
244 aac gcg ctg gtg acg ggc ggc agc cgc ggc att ggc gaa gcc act gcc
245 Asn Ala Leu Val Thr Gly Gly Ser Arg Gly Ile Gly Glu Ala Thr Ala
246 1
                                       10
248 att aag etc gee gag gag gge tac age gtc aeg att geg tet ege gge
249 Ile Lys Leu Ala Glu Glu Gly Tyr Ser Val Thr Ile Ala Ser Arg Gly
```

48

96

RAW SEQUENCE LISTING DATE: 08/29/2006
PATENT APPLICATION: US/10/573,130A TIME: 08:46:39

Input Set: A:\Substitute Sequence List-13111-00035-US.txt
Output Set: N:\CRF4\08292006\J573130A.raw

```
25
252 ctt aag cag ctc gag gct gtg aag gcc aaa cta ccc att gtg aag cag
253 Leu Lys Gln Leu Glu Ala Val Lys Ala Lys Leu Pro Ile Val Lys Gln
                               40
         35
256 gga cag gtt cac cac gtg tgg cag ctt gat ctc agt gat gtc gac gct
                                                                          192
257 Gly Gln Val His His Val Trp Gln Leu Asp Leu Ser Asp Val Asp Ala
       50
258
260 geg gee gee tte aaa ggg teg eeg eta eet gee age ege tae gae gtg
                                                                          240
261 Ala Ala Ala Phe Lys Gly Ser Pro Leu Pro Ala Ser Arg Tyr Asp Val
                                            75
264 ctc gtc agc aat gct ggc gtg gcc cag ttt agc ccg ttc atc gag cat
                                                                          288
265 Leu Val Ser Asn Ala Gly Val Ala Gln Phe Ser Pro Phe Ile Glu His
                                                            95
                                        90
                   85
266
268 geg aag cag gachtgg tog cag atg ott ged atd aat otg gog goa occ
                                                                          336
269 Ala Lys Gln Asp Trp Ser Gln Met Leu Ala Ile Asn Leu Ala Ala Pro
                                    105
               100
272 att geg etg gee cag aca ttt get aag gee att gge gae aag eeg ege
                                                                          384
273 Ile Ala Leu Ala Gln Thr Phe Ala Lys Ala Ile Gly Asp Lys Pro Arg
                                                    125
                                120
274
           115
276 aac aca ccg gcc cac att gtg ttt gtc tcg tcg aac gtc tcg ttg cga
                                                                          432
277 Asn Thr Pro Ala His Ile Val Phe Val Ser Ser Asn Val Ser Leu Arg
                                                140
                            135
280 ggc ttc ccg aac atc ggc gtc aac tcc atc acc ccc ggc a
                                                                          472
281 Gly Phe Pro Asn Ile Gly Val Asn Ser Ile Thr Pro Gly
                        150
282 145
285 <210> SEQ ID NO: 6
287 <211> LENGTH: 157
289 <212> TYPE: PRT
291 <213> ORGANISM: Candida magnoliae
294 <400> SEQUENCE: 6
296 Asn Ala Leu Val Thr Gly Gly Ser Arg Gly Ile Gly Glu Ala Thr Ala
                                        10
297 1
300 Ile Lys Leu Ala Glu Glu Gly Tyr Ser Val Thr Ile Ala Ser Arg Gly
                20
304 Leu Lys Gln Leu Glu Ala Val Lys Ala Lys Leu Pro Ile Val Lys Gln
                                40
            35
308 Gly Gln Val His His Val Trp Gln Leu Asp Leu Ser Asp Val Asp Ala
                                                 60
312 Ala Ala Ala Phe Lys Gly Ser Pro Leu Pro Ala Ser Arg Tyr Asp Val
                        70
316 Leu Val Ser Asn Ala Gly Val Ala Gln Phe Ser Pro Phe Ile Glu His
                                         90
317
320-Ala Lys Gln Asp Trp Ser Gln Met Leu Ala Ile Asn Leu Ala Ala Pro
                                                        110
                                    105
               100
 321
 324 Ile Ala Leu Ala Gln Thr Phe Ala Lys Ala Ile Gly Asp Lys Pro Arg
                                                     125
                                120
           115
 328 Asn Thr Pro Ala His Ile Val Phe Val Ser Ser Asn Val Ser Leu Arg
 329 130
                             135
 332 Gly Phe Pro Asn Ile Gly Val Asn Ser Ile Thr Pro Gly
```

RAW SEQUENCE LISTING DATE: 08/29/2006
PATENT APPLICATION: US/10/573,130A TIME: 08:46:39

Input Set : A:\Substitute Sequence List-13111-00035-US.txt
Output Set: N:\CRF4\08292006\J573130A.raw

```
150
                                             155
333 145
337 <210> SEQ ID NO: 7
338 <211> LENGTH: 27
339 <212> TYPE: DNA
340 <213> ORGANISM: Artificial sequence
342 <220> FEATURE:
343 <223> OTHER INFORMATION: Primer: Mke 338
345 <400> SEQUENCE: 7
                                                                             27
346 gggaattcca tatgtctaac cgtttgg
349 <210> SEQ ID NO: 8
350 <211> LENGTH: 28
351 <212> TYPE: DNA
352 <213> ORGANISM: Artificial sequence
354 <220> FEATURE:
355 <223> OTHER INFORMATION: Primer: Mke 339
357 <400> SEQUENCE: 8
                                                                             28
358 cgtagggaag cttattgagc agtgtagc
361 <210> SEQ ID NO: 9
362 <211> LENGTH: 28
363 <212> TYPE: DNA
364 <213> ORGANISM: Artificial sequence
366 <220> FEATURE:
367 <223> OTHER INFORMATION: Primer: Mke 366
369 <400> SEQUENCE: 9
                                                                             28
370 acgacgacga gcaacgcbct bgtbacgg
373 <210> SEQ ID NO: 10
374 <211> LENGTH: 28
375 <212> TYPE: DNA
376 <213> ORGANISM: Artificial sequence
378 <220> FEATURE:
379 <223> OTHER INFORMATION: Primer: Mke 367
381 <400> SEQUENCE: 10
                                                                             28
382 acgacgacgt cgaacgcbct bgtbacgg
385 <210> SEQ ID NO: 11
386 <211> LENGTH: 27
387 <212> TYPE: DNA
388 <213> ORGANISM: Artificial sequence
390 <220> FEATURE:
391 <223> OTHER INFORMATION: Primer: Mke 374
393 <400> SEQUENCE: 11
                                                                             27
394 gccggggttg atsswgttsa cgccgat
397 <210> SEQ ID NO: 12
398- <211> LENGTH: 10
399 <212> TYPE: PRT
400 <213> ORGANISM: Lactobacillus brevis
403 <220> FEATURE:
404 <221> NAME/KEY: MISC_FEATURE
405 <222> LOCATION: (1)..(10)
406 <223> OTHER INFORMATION: Fragment: C terminus
```

RAW SEQUENCE LISTING ERROR SUMMARY DATE: 08/29/2006 PATENT APPLICATION: US/10/573,130A TIME: 08:46:40

FYI

Input Set: A:\Substitute Sequence List-13111-00035-US.txt
Output Set: N:\CRF4\08292006\J573130A.raw

## lease Note:

se of n and/or Xaa have been detected in the Sequence Listing. Please review the equence Listing to ensure that a corresponding explanation is presented in the <220> o <223> fields of each sequence which presents at least one n or Xaa.

```
eq#:15; Xaa Pos. 60
eq#:16; Xaa Pos. 19,20
eq#:17; Xaa Pos. 12,13,14,15
eq#:18; Xaa Pos. 8
eg#:19; Xaa Pos. 9,11,12,13,14,15
eq#:22; Xaa Pos. 6,10,11,12,13,14,15
eq#:23; Xaa Pos. 6,12,15,16,17,18
eq#:24; Xaa Pos. 6,10,11,12,13,14,15,16,17
eq#:25; Xaa Pos. 16,17,18,19,20
eq#:26; Xaa Pos. 1,3
eq#:27; Xaa Pos. 9,13
eq#:28; Xaa Pos. 7,17,18,19,20
eq#:30; Xaa Pos. 1,10,11,12,13,14,15,16,17,18,19,20
eq#:32; Xaa Pos. 29,30
eq#:33; Xaa Pos. 5,6,7,8,9,10
eq#:34; Xaa Pos. 3,13,14,15,16,17,18,19,20
eq#:35; Xaa Pos. 11,12,13,14
eq#:36; Xaa Pos. 1
eq#:40; Xaa Pos. 1
eq#:41; Xaa Pos. 2,39,40
eq#:42; Xaa Pos. 37,38,39
eq#:43; Xaa Pos. 12,13,14,15
eq#:44; Xaa Pos. 13,14,15
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VERIFICATION SUMMARY
PATENT APPLICATION: US/10/573,130A

DATE: 08/29/2006 TIME: 08:46:40

Input Set : A:\Substitute Sequence List-13111-00035-US.txt
Output Set: N:\CRF4\08292006\J573130A.raw

```
1:496 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:15 after pos.:48
1:596 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:16 after pos.:16
1:617 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:17 after pos.:0
::634 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:18 after pos.:0
::705 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:19 after pos.:0
.:800 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:22 after pos.:0
.:897 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:23 after pos.:0
1:341 Repeated in SeqNo=23
.:923 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:24 after pos.:0
1:341 Repeated in SeqNo=24
1:944 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:25 after pos.:0
1:341 Repeated in SeqNo=25
,:970 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:26 after pos.:0
.:1057 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:27 after pos.:0
.:1154 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:28 after pos.:0
1:341 Repeated in SeqNo=28
::1320 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:30 after pos.:0
1:341 Repeated in SeqNo=30
::1554 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:32 after pos.:16
1571 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:33 after pos.:0
11632 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:34 after pos.:0
1:341 Repeated in SeqNo=34
 .:1653 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:35 after pos.:0
 ::1685 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:36 after pos.:0
1:1815 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:40 after pos.:0
1:1845 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:41 after pos.:0
1:341 Repeated in SeqNo=41
1:1987 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:42 after pos.:32
1:2004 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:43 after pos.:0
 ::2066 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:44 after pos.:0
```

<210>	15
<211>	60 PDM
<212>	PRT
<213>	Lactobacillus brewis
	VARIANT (47)(47) Amino acid is Ala or Lys  UNSURE (48)(48) Amino acid is Lys or Ala  VARIANT  VARIANT  VARIANT  Amino acid is Lys or Ala  VARIANT  VARIANT
	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
<220>	at exation of confirm
<221>	VARIANT HIA OF WELL
<222>	(47)(47)
<223>	Amino acid is Ala or Lys
\L_J>	
<220>	1/10 X a~
	Was the state of t
<221>	UNSURE CONTROLLED ON
<222>	(48)(48)
<223>	UNSURE (48)(48) Amino acid is Lys or Ala  Same evo  explain in
	Stocker 1
<220>	
<221>	VARIANT (53) (2267-62237
<222>	(53)(53)
<223>	Amino acid is Pro or Thr Same lund
1223-	(53)(53) Amino acid is Pro or Thr Sane euro  Setur
<220>	
<221>	VARIANT (59) (59)
	Jan 2
<222>	(05)11.105)
<223>	Amino acid is Phe, Val, Gly, or Asn
	·
<220>	
<221>	misc_feature
<222>	(60) . · (60) · · · · · · · · · · · · · · · · · · ·
<223>	Xaa is unknown
<400>	15
Ser As	n Arg Leu Asp Gly Lys Val Ala Ile Val Thr Gly Gly Thr Leu
1	5 10 15
•	-
C1 T3	e Gly Leu Ala Ile Ala Thr Lys Phe Val Glu Glu Gly Ala Lys
GIY II	
	20 25 30
	The state of the s
Val Me	t lle Thr Gly Arg His Ser Asp Val Gly Glu Lys Ala (Ala Lys)
	35 40 45 . · · · · ·
Ser Va	al Gly Thf Pro Asp Gln Ile Gln Phe Phe Xaa
50	al Gly The Pro Asp Gln Ile Gln Phe Phe Xaa 60
-	
	a // A
	. I. Ih and
_	In sequence !
	1 1 sant agreer
	1 / tuple   eve
1	Lese Joyle )
/	Lese types I ever appear in Sequence 16 and
	- u

subsequent